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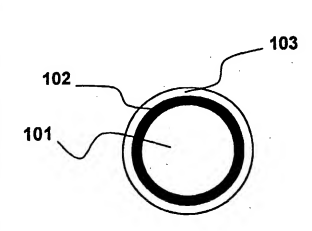
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(54) Title: OPTICAL FIBER WITH EPOXIDIZED POLYOLEFIN BASED COATING



(57) Abstract: Optical fiber comprising at least one epoxidized polyolefin based polymer coating, characterized in that said coating is formed from a crosslinkable composition comprising: (a) at least one epoxidized polydiene oligomer having a first and a second end, said oligomer comprising at least one hydrocarbon chain that is substantially free of ethylenic double bonds, at least one epoxide group at said first end and at least one reactive functional group at said second end; b) at least one hydrogenated polydiene oligomer comprising at least one reactive functional group capable of reacting with said epoxide groups; (c) at least one photo-initiator. Preferably, said coating is a primary coating, preferably coated with a secondary coating.

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### "OPTICAL FIBER WITH EPOXIDIZED POLYOLEFIN BASED COATING"

The present invention relates to an optical fiber comprising at least one epoxidized polyolefin based coating layer, and to a crosslinkable composition which can be applied as said coating.

More particularly, the present invention relates to an optical fiber comprising at least one epoxidized polyolefin based primary coating layer and at least one secondary coating layer deposited around said primary coating, and to a crosslinkable composition which can be applied as said primary coating.

Optical fibers commonly consist of a glass portion (typically with a diameter of about 125  $\mu m$ ), inside which the transmitted optical signal is confined, and of 15 a coating, typically polymeric, arranged around the glass portion for substantially protective purposes. This protective coating typically comprises a first coating layer positioned directly onto the surface, known as the "primary coating" or "primary" for 20 short, typically having a thickness of between about 25  $\mu\mathrm{m}$  and about 35  $\mu\mathrm{m}$ . In turn, this primary coating is generally covered with a second coating Tayer, known as "secondary coating" or "secondary" typically having a thickness of between about 10  $\mu\mathrm{m}$  and 25 about 30  $\mu$ m.

polymer coatings may ' be obtained compositions comprising oligomers and monomers that are generally crosslinked by means of UV irradiation in the presence of a suitable photo-initiator. The two coatings described above differ, inter alia, in terms of the modulus of elasticity of the crosslinked material. As a matter of fact, whereas the material which forms the primary coating is a relatively soft material, with a 35 relatively low modulus elasticity of at temperature, the material which forms the secondary

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coating is relatively harder, having higher modulus of elasticity values at room temperature. The combination of said two layers of coating ensures adequate mechanical protection for the optical fiber.

The optical fiber thus composed usually has a total diameter of about 250  $\mu m$ . However, for particular applications, this total diameter may also be smaller; in this case, a coating of reduced thickness is generally applied.

The crosslinking of the abovementioned compositions, depending on the reactive groups present in the compounds (oligomers and monomers) to be crosslinked, may take place, for example, via a free-radical or cationic route. Typically, the crosslinking of compounds comprising epoxide groups takes place cationically.

For example, patent application EP 124 057 describes a cationically crosslinkable liquid composition comprising a polyepoxide, a polysiloxane bearing a plurality of hydroxyalkyl groups in the molecule, and a photo-initiator and/or a photo-sensitizer. According to the assertions made in the application, said composition is capable of providing a coating for optical fibers what is capable of maintaining low modulus of elasticity values at low temperatures (-60°C) so as to avoid the phenomenon known as "microbending", with consequent attenuation of the transmitted signal.

Patent application EP 533 397 describes an optical fiber with a coating which includes at least one layer comprising a cationically crosslinkable composition. resin containing comprises а composition 30 Said cationically crosslinkable end groups, diluent containing cationically crosslinkable end groups, and a photo-initiator. Resins that are useful for this purpose may be selected from vinyl ether resins and epoxy the According to the assertions made resins. 35 application, said cationic crosslinking leads to

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formation of an acid medium in contact with the glass portion of the optical fiber, making it possible to obtain an optical fiber with improved mechanical strength.

5 Patent US 6 042 943 describes а radiationcrosslinkable composition which may be used as primary coating for an optical fiber, comprising: (a) a compound comprising (i) a saturated aliphatic chain, and (ii) at least one epoxide group at one end and at least one reactive function, which may be selected from acrylates, 10 vinyl ethers, hydroxyls, of combinations thereof, at the other end; (b) a blend of acrylate-type monomers comprising (iii) a first monomer containing an acrylate group, and (iv) a second monomer containing at least two 15 acrylate groups; and (c) a photo-initiator. According to further embodiment, (b) is a blend of monomers comprising (iii) a first monomer containing an acrylate group or a vinyl ether group, and (iv) a second monomer containing at least two functional groups which may be selected from acrylates, epoxides, vinyl ethers and 20 hydroxyls. According to a further embodiment, (b) is a blend of monomers of vinyl ether type comprising (iii) a first monomer containing a vinyl ether group, and (iv) a second monomer containing at least two vinyl ether 25 groups. According to a further embodiment, (b) is a monoacrylate residue containing from 6 to 20 carbon atoms.

Patent US 5 993 965 describes a fiber with a coating based on a hydrophobic material derived from the photopolymerization of a composition comprising at least one epoxidized polydiene oligomer, at least one photoinitiator and, optionally, a reactive diluent monomeric type. According to the assertions made in the patent, the fibers thus coated are said to have improved mechanical behaviour. In particular, said coating is used in optical fibers.

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As observed by the Applicant, the use of reactive diluents of monomeric type, that are generally required to obtain compositions whose viscosity allows them to be applied to optical fibers at room temperature, may present some drawbacks. For example, the relatively low molecular weight of these monomer components connected with a relatively high volatility, with a consequent contamination of other materials and/or risks to the environment and to the health of the workers. In addition, following the crosslinking by UV irradiation, residues of unreacted components may remain in the final resin. The presence of these unreacted monomer residues within the polymer network may result in unwanted phenomena of extraction by water and/or waterblocking fillers commonly used in optical cables to prevent or limit the entry of water into the structure of the cable. This extraction entails a worsening in mechanical properties and may also result in the initiation of the phenomenon of delamination of the fiber, i.e. detachment of the polymer coating from the glass portion of the fiber, with possible generation of the phenomenon known as "microbending".

The Applicant has now found that the use, of a hydrogenated polydiene oligomer comprising at least one 25 reactive function, preferably at least one reactive end function, makes it possible to obtain compositions with an acceptable viscosity at room temperature, with little or no use of conventional diluent monomers. The use of a composition according to the invention makes it possible to obtain a polymer coating for an optical fiber, in 30 particular a primary coating, with improved properties modulus example, relatively low as, for elasticity values at the normal working temperatures of said fiber, in particular at low temperatures. Said 35 compositions show reduced toxicity by virtue of the lower volatility of the components and, thus, fewer

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risks not only as regards the contamination of other materials, but also as regards the environment and the health of the workers. In addition, said compositions show improved behaviour both in the presence of water and in the presence of waterblocking fillers.

The Applicant has also observed that, while conventional polymer coatings are applied at temperature, the use of the abovementioned oligomer instead of the abovementioned reactive diluents of the monomeric type makes it possible to work at higher 10 application temperatures. The possibility of working at higher temperatures without the risk of volatilization of the low molecular weight components makes it possible also to use compositions that, at room temperature, have a viscosity that is higher than those normally used, to 15 increase the crosslinking rate and to avoid a further crosslinking treatment ("post-curing") of the alreadycoated optical fiber. Higher application temperatures allows also to avoid cooling the fiber temperature before the application of the coating. 20

In particular, the abovementioned oligomer is advantageously used as a blend with an epoxidized polydrene oligomer comprising at least one hydrocarbon chain that is substantially free of ethylenic double bonds.

According to a first aspect, the present invention thus relates to an optical fiber comprising at least one epoxidized polyolefin based polymer coating, characterized in that said coating is formed from a crosslinkable composition comprising:

(a) at least one epoxidized polydiene oligomer having a first and a second end, said oligomer comprising at least one hydrocarbon chain that is substantially free of ethylenic double bonds, at least one epoxide group at said first end and at least one reactive functional group at said second end;

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- (b) at least one hydrogenated polydiene oligomer comprising at least one reactive functional group capable of reacting with said epoxide groups;
- (c) at least one photo-initiator.
- According to a preferred embodiment of the present invention, said composition optionally comprises at least one adhesion promoter (d).

According to a further preferred embodiment, said polymer coating is a primary coating, preferably coated with a secondary coating.

- relates to a crosslinkable composition comprising:
- (a) at least one epoxidized polydiene oligomer having a first and a second end, said oligomer comprising at least one hydrocarbon chain that is substantially free of ethylenic double bonds, at least one epoxide group at said first end and at least one reactive functional group at said second end;
- (b) at least one hydrogenated polydiene oligomer 20 comprising at least one reactive functional group capable of reacting with said epoxide groups;
  - (c) at least one photo-initiator.

invention, said composition optionally comprises at least one adhesion promoter (d).

According to a further embodiment of the present invention, said composition optionally comprises at least one reactive diluent monomer (e).

According to a preferred embodiment, said 30 crosslinkable composition has a modulus of elasticity, at room temperature, of less than about 4 MPa, preferably between 1 MPa and 3 MPa.

According to a further preferred embodiment, said crosslinkable composition has a modulus of elasticity, at -40°C, of between 5 MPa and 350 MPa, preferably between 10 MPa and 50 MPa.

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Said modulus of elasticity is measured using DMTA (Dynamic Mechanical Thermal apparatus Analyser from Reometrics Inc.), in traction, at a frequency of 1 Hz and at a heating rate of 2°C/min.: further details regarding the analysis method will be described in the examples given hereinbelow.

According to a further aspect, the present invention relates to method for a applying an epoxidized polyolefin based polymer coating to an optical fiber, which comprises:

- drawing a glass preform placed in a suitable furnace;
- cooling the fiber leaving the furnace;
- applying said coating;
- 15 crosslinking said coating; characterized in that the application of said coating layer is carried out at a temperature of not less than 60°C, preferably between 80°C and 120°C.

Preferably, the abovementioned coating is formed from a crosslinkable composition comprising: 20

- (a) at least one epoxidized polydiene oligomer having a first and a second end, said oligomer comprising at least one hydrocarbon chain that is substantiallyfree of ethylenic double bonds, at least one epoxide group at said first end and at least one reactive functional group at said second end;
  - (b) at least one hydrogenated polydiene oligomer comprising at least one reactive functional group capable of reacting with said epoxide groups;
  - (c) at least one photo-initiator. 30 Generally, said epoxidized polydiene oligomer (referred to hereinbelow for simplicity as "epoxidized compound (a)") is, at room temperature, in the form of a viscous liquid.
  - 35 The epoxidized compound (a) is generally prepared by anionic (co)polymerization of conjugated diene monomers

to give a polydiene according to known techniques as described, for example, in patents US 5 247 026, US 5 536 772, US 5 264 480, US 6 042 943 and patent application EP 516 203.

For example, said (co)polymerization may be carried 5 out in bulk, in solution or in emulsion. In general, in the solution (co)polymerization, an initiator selected, for example, from metals belonging to group IA of the Table of the Elements, or alkyl, amide, Periodic biphenyl, anthracenyl derivatives silanol. naphthyl, 10 thereof, is used, and the polydiene is obtained by (co)polymerizing, simultaneously or sequentially, the The (co)polymerization monomers. diene conjugated reaction is generally carried out at a temperature of between about -150°C and about 300°C, preferably between 15 0°C and 100°C, in a suitable solvent. Preferably, the (co)polymerization initiator is selected from organic for example, compounds of alkali metals such as, organolithium compounds represented by the following 20 general formula:

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in which R represents an aliphatic, cycloaliphatic or aromatic: hydrocarbon a said a aromatic phydrocarbon with alkyl groups substituted optionally being containing from 1 to 20 carbon atoms, and n is an 25 integer between 1 and 4. The polydiene thus obtained may the residual reacting be functionalized by organometallic groups derived from said initiator by reaction with suitable terminating agents such as, for example, low molecular weight alkylene oxides in the 30 presence of small amounts of aliphatic tertiary amines  $N, N, N^1, N^1$ -tetramethyleneexample, for such as, ethylenediamine.

The polydiene thus obtained is then hydrogenated and epoxidized according to known techniques as described, for example, in patent US 4 879 349. The process

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described in said patent involves the (co)polymerization of substituted or unsubstituted conjugated dienes and the subsequent hydrogenation of the copolymer, working under conditions such that the unsubstituted ethylenic unsaturations present in the copolymer are selectively hydrogenated, while the substituted ethylenic unsaturations present in said copolymer remain substantially non-hydrogenated. The · partially hydrogenated polydiene thus obtained is subsequently epoxidized according to known techniques, for example by reaction with an organic peracid (for example peracetic acid or perbenzoic acid).

The hydrogenation and the epoxidation may be carried out in any order. Preferably, the polydiene is first hydrogenated and then epoxidized.

According to a preferred embodiment, the epoxidized compound (a) is obtained by anionic (co)polymerization of conjugated diene monomers containing from 4 to 24, preferably from 4 to 12 carbon atoms selected, 20 example, from: isoprene, 1,3-butadiene, 2-ethyl-1,3butadiene, 2-buty1-1,3-butadiene, 2-penty1-1,3butadiene, 2-hexyl-1,3-butadiene, 2-heptyl-1.3butadiene, 2-octyl-1,3-butadiene, 2-nonyl-1,3-butadiene, 2-decyl-1,3-butadiene, 2-dodecy1-1,3-butadiene, 2tetradecy1-1,3-butadiene, 2-hexadecy1-1,3-butadiene, 25 isoamyl-1,3-butadiene, 2-phenyl-1,3-butadiene, 2-methyl-1,3-pentadiene, 2-methyl-1,3-hexadiene, 2-methy1-1,3heptadiene, 2-methyl-1,3-octadiene, 2-methy1-6methylene-2,7-octadiene, and mixtures thereof. Disubstituted conjugated diene monomers such as, for 30 2,3-dimethy1-1,3-butadiene, 2,3-dimethy1-1,3example, pentadiene, 2,3-dimethy1-1,3-hexadiene, 2,3-dimethy1-1,3-heptadiene, 2,3-dimethyl-1,3-octadiene, or mixtures thereof, may also be used. Alternatively, difluorinated conjugated diene monomers such as, for example, 2,3-35 difluoro-1,3-butadiene, 2,3-difluoro-1,3-pentadiene,

2,3-difluoro-1,3-hexadiene, 2,3-difluoro-1,3-octadiene, or mixtures thereof, may be used. 1,3-butadiene and isoprene are preferred.

The conjugated diene monomers may optionally be other ethylenically 5 with unsaturated copolymerized monomers such as, for example:  $\alpha$ -olefins containing from 2 to 12 carbon atoms (for example ethylene, propylene, 1-butene), monovinylarenes containing from 8 carbon atoms (for example styrene, 1-vinylnaphthalene, 10 3-methylstyrene), vinyl esters in which the ester group contains from 2 to 8 carbon atoms (for example vinyl acetate, vinyl propionate, vinyl butanoate), acrylates and alkyl methacrylates in which the alkyl contains from 1 to 8 carbon atoms (for example ethyl 15 acrylate, methyl acrylate, t-butyl acrylate, n-butyl acrylate), acrylonitrile, or mixtures thereof.

Preferably, the epoxidized compound (a) is prepared by sequential anionic (co)polymerization working as follows:

- 20 (a) anionic polymerization of a first conjugated diene monomer, in particular isoprene;
- (b) anionic copolymerization of the polymer obtained in stage (a) with a second conjugated diene monomer, in particular 1,3-butadiene, thus obtaining a block copolymer;
  - (c) functionalization of the block copolymer obtained in stage (b) with a suitable terminating agent, in particular ethylene oxide;
- (d) subsequent selective hydrogenation and epoxidation of the functionalized block copolymer obtained in stage (c).

According to a preferred embodiment, the epoxidized compound (a) is a diblock copolymer comprising a first block comprising at least one epoxide group, obtained by the polymerization of a first conjugated diene monomer,

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in particular isoprene, which is subsequently epoxidized, and a second block formed from a hydrocarbon chain that is substantially free of ethylenic double bonds, obtained by the polymerization of a second conjugated diene monomer, in particular 1,3-butadiene, subsequently terminated with at least one reactive functional group and hydrogenated. In this case, the hydrocarbon chain that is substantially free ethylenic double bonds is a poly(ethylene/butylene) chain.

Groups selected, for example, from aliphatic groups, cycloaliphatic groups, aryl groups, combinations thereof, may optionally be inserted into the hydrocarbon chain that is substantially free of ethylenic double bonds. Said groups are not incorporated into the main hydrocarbon chain that is substantially free of ethylenic double bonds, but are present in a side chain. In this case, in the epoxidized compound (a), the hydrocarbon chain that is substantially free of ethylenic double bonds may be, for example, poly(ethylene/butylene/styrene).

Preferably, said hydrocarbon chain that is substantially free of ethylenic double bonds has an average (number-average) molecular weight, which may be determined for example by gel permeation chromatography (GPC), of between 2,000 daltons and 10,000 daltons, preferably between 3,000 daltons and 6,000 daltons.

As stated above, the epoxidized compound (a) contains at least one epoxide group at the first end. The number of epoxide groups present in the epoxidized compound (a) may vary according to the epoxidation process used.

According to a preferred embodiment, from 5 to 15 epoxide groups, preferably from 9 to 11 epoxide groups are present in the epoxidized compound (a), at the first end.

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stated above, the epoxidized compound As contains at least one reactive functional group at the second end. The expression "reactive functional group" means a group that can react with compounds such as, for 5 example, acrylates, vinyl ethers, epoxides, alcohols or isocyanates. Said reactive functional group may be selected, for example, from hydroxyl, acrylate, epoxy, vinyl ether, mercaptan. When two or more reactive functional groups are present, said groups may be identical to or different from each other.

According to a preferred embodiment, in the epoxidized compound (a), the reactive functional group present at the second end is a hydroxyl group.

When the reactive functional group is a hydroxyl, 15 said group may be converted into other reactive functional groups using techniques known in the art.

The epoxidized compound (a) may be prepared in various forms according to the technique used.

According to a preferred embodiment, the epoxidized compound (a) is a linear, star or radial polymer. The epoxidized compound (a) is preferably a linear polymer.

According to a further preferred embodiment, the epexidized compound (a) has an average (number-average) molecular weight of between 3,000 daltons and 15,000 daltons, preferably between 5,000 daltons and 7,000 daltons. Said average molecular weight may be determined as described above.

According to a further preferred embodiment, the epoxidized compound (a) has a viscosity, measured at 30°C, of less than 1,000 poise, preferably less than 600 30 poise, up poise. Said viscosity to 100 determined, for example, using viscometer a of Brookfield type, model DV-III, equipped configuration 29.

35 Epoxidized compounds (a) which may be used in the present invention are commercially available, for

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example, under the brand name Kraton Liquid<sup>TM</sup> Polymer from Kraton Polymer. Kraton Liquid<sup>TM</sup> EKP-207 is particularly preferred.

The substantial absence of ethylenic double bonds in the hydrocarbon chain of the epoxidized compound (a) is particularly preferred for the purposes of the present invention, since their presence can cause degradation phenomena (thermal or oxidative degradation degradation by exposure to ultraviolet light) of said compound. Any degradation might in its turn involve a 10 worsening in the mechanical properties of the polymer coating of the optical fiber. In addition, unwanted phenomena of coloration of said coating might be encountered.

15 According to preferred embodiment, a the hydrogenated polydiene oligomer (b) (referred hereinbelow for simplicity as "hydrogenated compound (b)") generally comprises a base polymer structure of synthetic or natural origin, which is derived from the (co)polymerization of one or more conjugated diene 20 monomers, optionally copolymerized with other ethylenically unsaturated monomers.

Conjugated diene monomers that are particularly preferred for the purposes of the present invention are those containing from 4 to 24 carbon atoms, preferably from 4 to 12 carbon atoms, selected, for example, from: 1,3-butadiene, isoprene, piperylene, methylpentadiene, phenylbutadiene, 3,4-dimethyl-1,3-hexadiene, 4,5-diethyl-1,3-octadiene, or mixtures thereof. 1,3-butadiene and isoprene are particularly preferred.

Ethylenically unsaturated monomers that are particularly preferred according to the present invention are, for example:  $\alpha$ -olefins containing from 2 to 12 carbon atoms (for example ethylene, propylene, 1-butene), monovinylarenes containing from 8 to 20 carbon

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atoms (for example styrene, 1-vinylnaphthalene, 3-methylstyrene), vinyl esters in which the ester group contains from 2 to 8 carbon atoms (for example vinyl acetate, vinyl propionate, vinyl butanoate), alkyl acrylates and alkyl methacrylates in which the alkyl contains from 1 to 8 carbon atoms (for example ethyl acrylate, methyl acrylate, methyl methacrylate, t-butyl acrylate, n-butyl acrylate), acrylonitrile, or mixtures thereof.

10 According to preferred embodiment, in the hydrogenated compound (b), the reactive functional group may be selected, for example, from: hydroxyl, acrylate, epoxy, vinyl ether, mercaptan. When two or more reactive functional groups are present, said groups may be identical to or different from each other. Preferably, 15 said reactive functional group is a hydroxyl group, more preferably it is a hydroxyl group in an end position. Preferably, the hydrogenated compound (b) has a hydroxyl functionality of between about 0.5 and about 2.6. Said 20 hydroxyl functionality may be determined, for example, according to ASTM standard E222-00.

According to a further preferred embodiment, the hydrogenated compound (b) has a viscosity, measured at 30°C, of between about 10 poise and about 1,000 poise, more preferably between about 20 poise and about 400 poise. Said viscosity may be determined as described above.

According to a further preferred embodiment, the hydrogenated compound (b) has an average (number-average) molecular weight of between about 500 daltons and about 20,000 daltons, more preferably between about 2,000 daltons and about 10,000 daltons. Said average molecular weight may be determined as described above.

According to a further preferred embodiment, the 35 hydrogenated compound (b) containing at least one hydroxyl end function has a hydroxyl-equivalent weight

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of between about 250 and about 20,000, preferably between about 500 and about 10,000. Said hydroxylequivalent weight may be determined, for example, according to ISO standard 3001:1999.

- Hydrogenated compounds (b) that may be used in the 5 present invention are commercially available, example, under the brand name Kraton Liquid $^{\text{TM}}$  Polymer from Kraton Polymer. Kraton Liquid $^{\text{TM}}$  L-1203 Polymer and L-2203 Polymer are particularly preferred.
- 10 hydrogenated compound (b) The may be prepared according to known techniques. For example, the base polymer may be prepared by (co)polymerization of the corresponding monomers in emulsion, in suspension or in solution. In particular, the base polymers obtained by 15 anionic polymerization in the presence of an organometallic initiator (in particular an organolithium initiator) may be functionalized by reacting the residual organometallic groups derived from initiator by reaction with suitable terminating agents such as, for example, alkylene oxides or low molecular 20 weight epoxides in the presence of small amounts of aliphatic tertiary amines such as, for example,  $N, N, N^1, N^1$ -tetramethyleneethylenediamine. Further details regarding the preparation of the hydrogenated polydiene

oligomers described above are given, for example, in 25 patents US 4 039 593 and US 5 916 941. According to a preferred embodiment, the photo-

initiator (c) may be selected from salts that are capable of forming strong acids when subjected to UV irradiation so as to initiate the cationic crosslinking.

Specific examples of photo-initiators (c) which may be used in the present invention are: hexafluorophosphorus triarylsulphonium salts. hexafluoroantimony triarylsulphonium salts, (tolylcumyl) tetrakis(pentafluorophenyl)iodonium salts, diaryliodonium hexafluoroantimonate

salts,

or

mixtures

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thereof.

stated above, the crosslinkable composition according to the present invention may optionally comprise a reactive diluent monomer (e).

5 According to a preferred embodiment, said reactive diluent monomer (e) may be selected from vinyl ethers such as, for example, n-butyl vinyl ether, n-dodecyl vinyl ether, or mixtures thereof.

For the purposes of limiting the problems outlined 10 above relating to the use of reactive diluent monomers, ....it is preferable for the amount of reactive diluent monomer added to the crosslinkable composition to be not greater than 20 parts by weight, preferably between 0 parts by weight and 10 parts by weight relative to 100 15 parts by weight of (a) + (b).

the crosslinkable composition stated above, the present invention may optionally according to least one adhesion promoter (d). comprise at provides increased adhesion adhesion promoter (d) between the glass fiber and the primary coating. As observed by the Applicant, while the adhesion between the crosslinked composition and the glass is generally re-acceptable, in particular on the freshly manufactured this adhesion may nevertheless optical fiber, impaired upon ageing, with possible undesirable reduction of said adhesion strength. The use of a suitable adhesion promoter thus allows to maintain the value of adhesion strength at an acceptable value, also upon ageing of the optical fiber.

30 Said adhesion promoter (d) is preferably an organofunctional silane.

For the purpose of the present description and the claims, the term "organo-functional silane" is intended to indicate a silyl compound with functional groups that facilitate the chemical or physical bonding between the glass surface and the silane, which ultimately results

in increased or enhanced adhesion between the primary coating and the glass fiber.

Specific examples of organo-functional silanes that used in the present invention are: octyltriethoxysilane, methyltriethoxysilane, methyltri-5 methoxysilane, tris(3-trimethoxysilylpropyl)isocyanurate, vinyltriethoxysilane, vinyltrimethoxysilane, vinyl-tris(2-methoxyethoxy)silane, vinylmethyldimethoxysilane, gamma-methacryloxypropyltrimethoxysilane, beta(3,4-epoxycyclohexyl)ethyltrimethoxy-silane, 10 gamma-glycidoxypropyltrimethoxysilane, gamma-mercaptopropyltrimethoxysilane, organo-modified polydimethylsiloxane, gamma-ureidopropyltrialkoxysilane, gamma-ureidopropyltrimethoxysilane, gamma-isocyanate-15 propyltriethoxysilane, or mixtures thereof. gamma-Glycidoxypropyltrimethoxysilane, beta(3,4-epoxycycloexhyl)ethyltrimethoxysilane and gamma-mercaptopropyltrimethoxysilane, are particularly preferred. gamma-Mercaptopropyltrimethoxysilane is more particularly 20 preferred.

Other examples of organo-functional silanes that may be used in the present invention may be identified, for example, by the following structural formula (I):

 $(R)_3Si-C_nH_{2n}-X \qquad (I)$ 

in which the groups R, which may be identical to or 25 different from each other, are chosen from: alkyl, alkoxy or aryloxy groups or from halogen atoms, condition that at least one of the groups R is an alkoxy or aryloxy group; n is an integer between 1 and 6 30 inclusive; X is group selected from: nitrous, mercapto, epoxide, vinyl, imido, chloro,  $\text{-(S)}_{m}C_{n}H_{2n}\text{-Si-(R)}_{3}$  in which m and n are integers between 1 and 6 inclusive and the groups R are defined as above. Among these, bis(3-trimethoxysilylpropyl)disulfane and 35 bis(3-tri-ethoxysilylpropyl)disulfane, are particularly preferred.

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Adhesion promoters (d) that may be used in the present invention are commercially available, for example, under the brand name Silquest® A-187 and Silquest® A-186 from OSi Specialties, Dynasylan® MTMO and Si® 266 from Degussa-Hüls.

The adhesion promoter is preferably added to the crosslinkable composition in an amount of from 0.1 parts by weight to 2.5 parts by weight, more preferably of from 0.3 parts by weight to 1.5 parts by weight relative to 100 parts of (a) + (b).

Conventional additives may be added for the purpose improving the fundamental characteristics of the abovementioned composition. For example, solvents, surfactants capable plasticizers, of improving wettability ("wetting") of the coating on the glass 15 portion of the optical fiber, devolatilizing agents, rheological agents, antioxidants, UV stabilizers capable of not interfering with the crosslinking operations may be added.

- 20 According to one preferred embodiment, the crosslinkable composition comprises:
- (a) about 20-80 parts by weight of at least one epoxidized polydiens oligomer having a first and a second end comprising at least one hydrocarbon chain that is substantially free of ethylenic double bonds, at least one epoxide group at said first end and at least one reactive functional group at said second end;
- (b) about 20-80 parts by weight of at least one 30 hydrogenated polydiene oligomer comprising at least one reactive functional group capable of reacting with said epoxide groups;
  - (c) about 0.05-5 parts by weight relative to 100 parts of (a) + (b) of a photo-initiator.
- According to a further preferred embodiment, said crosslinkable composition further comprises about 0.1

parts by weight to 2.5 parts by weight relative to 100 parts of (a) + (b) of an adhesion promoter (d).

As stated above, the abovementioned crosslinkable composition is particularly useful as a primary coating for an optical fiber. Said primary coating is then coated with a secondary coating that is compatible therewith. For example, a secondary coating formed from a crosslinkable composition comprising an epoxidized polydiene oligomer, a reactive diluent monomer and at least one photo-initiator may be used.

The epoxidized polydiene oligomer generally represents from 30% to 70% by weight of the secondary coating composition. The epoxidized polymer is preferably a hydrocarbon polyol such as, for example, partially hydrogenated and epoxidized polybutadiene containing two hydroxyl end groups and internal epoxide groups along the chain.

reactive The diluent monomer present in composition of the secondary coating is generally used in an amount of up to 400 parts per 100 parts by weight 20 epoxidized polymer, preferably in an amount of between 40 and 200 parts by weight of. epoxidized polymer. Reactive diluent monomers of epoxide type that may advantageously be used are, for example: 3,4-epoxy-25 cyclohexylmethyl-3,4-epoxycyclohexane carboxylate, limonene epoxide, cyclohexene epoxide, 1,2epoxydodecane. Reactive diluent monomers of vinyl ether type that may advantageously be used are, for example: triethylene glycol divinyl ether, 1,4-butanediol 30 monovinyl 1,4-bis(vinyloxymethyl)cyclohexane. ether, Reactive diluent monomers of oxetane type such as, for example: trimethylene oxide, 3,3-dimethyloxetane, 3,3dichloromethyloxetane, 3-ethyl-3-phenoxymethyloxetane, bis(3-ethyl-3-methyloxy)butane, or mixtures thereof, may 35 also advantageously be used.

The photo-initiator present in the composition of

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the secondary coating is generally used in an amount of up to 10 parts per 100 parts by weight of epoxidized polymer, preferably in an amount of between 0.01 and 10 parts. Photo-initiators that may advantageously be used are: hexafluorophosphorus triarylsulphonium salts, hexafluoroantimony triarylsulphonium salts, (tolylcumyl)tetrakis(pentafluorophenyl)iodonium salts, diaryliodonium hexafluoroantimonate salts, or mixtures thereof.

Further additives may be added for the purpose of 10 improving the fundamental characteristics of the composition of the secondary coating. For example, solvents, levelling agents, surface tension modifiers, modifiers, plasticizers, attrition coefficient surfactants, devolatilizing agents, rheological agents, 15 of antioxidants, and UV stabilizers capable interfering with the crosslinking operations may be added.

Said composition of the secondary coating preferably
has a modulus of elasticity at room temperature of less
than 2,500 MPa, preferably between about 300 MPa and
about 2,000 MPa. Said modulus of elasticity may be
determined by means of a DMPA analyser as described
above.

25 Compositions of the type described above which may be used as a secondary coating according to the present invention are described, for example, in patent US 5 993 965.

The present invention may be understood more clearly with reference to the following attached figures:

Figure 1: is a cross section of an optical fiber according to the invention;

Figure 2: is the general scheme of a system (spinning tower) for producing an optical fiber according to the invention.

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Figure 1 shows an optical fiber according to the

present invention, comprising a glass portion (101) covered with a primary coating (102) which may be produced according to the present invention, and which in its turn is covered with a secondary coating (103).

An optical fiber according to the present invention may be produced according to the usual spinning techniques, using, for example, a system such as the one schematically illustrated in Figure 2.

This system, commonly known as a "drawing tower", typically comprises a furnace (302) 10 inside which is placed a glass optical preform to be drawn. The bottom part of said preform is heated to the softening point and drawn into an optical fiber (301). The fiber is then cooled, preferably to a temperature of not less than 60°C, preferably in a suitable cooling tube (303) of the 15 type described, for example, in patent application WO 99/26891, and passed through a diameter measurement device (304). This device is connected by means of a microprocessor (313) to a pulley (310) which regulates the spinning speed; in the event of any variation in the 20 diameter of the fiber, the microprocessor (313) acts to regulate the rotational speed of the pulley (310), so as to keep the diameter of the optical fiber constant. the fiber passes through a primary coating applicator (305), containing the coating composition in 25 liquid form, and is covered with this composition to a thickness of about 25  $\mu$ m-35  $\mu$ m. As stated above, the application of the primary coating is preferably carried out at a temperature of at least 60°C, preferably between 80°C and 100°C. The coated fiber is then passed 30 through a UV oven (or a series of ovens) (306) in which the primary coating is crosslinked. The fiber covered with the crosslinked primary coating is then passed through a second applicator (307), in which it is coated with the secondary coating and then crosslinked in the relative VU oven (or series of ovens) (308).

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Alternatively, the application of the secondary coating may be carried out directly on the primary coating before the latter has been crosslinked, according to the "wet-on-wet" technique. this In case, a applicator is used, which allows the sequential application of the two coating layers, for example, of the type described in patent US 4 474 830. The fiber thus covered is then crosslinked using one or more UV ovens similar to those used to crosslink the individual coatings.

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Subsequently to the coating and to the crosslinking of this coating, the fiber may optionally be made to pass through a device capable of giving a predetermined torsion to this fiber, for example of the type described 15 in international patent application WO 99/67180, for the purpose of reducing the PMD ("Polarization Dispersion") value of this fiber. The pulley placed downstream of the devices illustrated previously controls the spinning speed of the fiber. After this drawing pulley, the fiber passes through a device (311) 20 capable of controlling the tension of the fiber, of the type described, for example, in patent application EP 1 112 979, and is finally collected on a reel (312).

An optical fiber thus produced may be used in the production of optical cables. The fiber may be used either as such or in the form of ribbons comprising several fibers combined together by means of a common coating.

Although the present invention has been described 30 with particular reference to a primary coating which is in turn coated with a secondary coating, according to the abovementioned description it is apparent to those skilled in the art that a crosslinkable formulation according to the present invention may be suitably formulated so as to be used as a secondary coating, or as a single coating for an optical fiber.

The present invention will be further illustrated hereinbelow by means of a number of implementation examples that are provided purely as a guide and are non-limiting on the invention.

#### EXAMPLES 1-5

## Preparation of compositions for primary coating

Compositions for the primary coating according to the invention were prepared: the amounts of the components (parts by weight except where otherwise 10 mentioned) are given in Table 1. The set that I have been been as well as I have

TABLE 1

	COMPOSITIONS				
COMPONENTS	1 (*)	2 (*)	3	4	5
Poly Bd® 605	76	-			-
Kraton® Liquid L-207	-	80	50	50	50`
Kraton® Liquid L-1203	- 200 - 4 2 D	- -14,71,81	50	-	30
Kraton® Liquid L-2203	-	_	-	50	20
Rapicure® HBVE	24	-		,	-
Rapicure® CHVE	-	20	-	-	
UVI® 6974  (*): comparative	1	1	1	1	1

<sup>(\*):</sup> comparative

Poly Bd® 605: epoxidized polybutadiene sold by Elf 15 Atochem;

Kraton Liquid  $^{\text{TM}}$  EKP-207: linear oligomer containing a poly(ethylene/butylene) aliphatic chain, a hydroxyl group at one end and epoxide groups at the other end,

sold by Kraton Polymer;

Kraton Liquid<sup>™</sup> L-1203: hydroxy-terminated hydrogenated polydiene oligomer sold by Kraton Polymer;

Kraton Liquid<sup>™</sup> L-2203: dihydroxy-terminated hydrogenated polydiene oligomer sold by Kraton Polymer;

Rapicure® HBVE: 4-hydroxybutyl vinyl ether sold by ISP; Rapicure® CHVE: cyclohexane dimethanol vinyl ether sold by ISP;

UVI® 6974: hexafluoroantimony triarylsulphonium salt, a photo-initiator sold by Union Carbide, as a 50% dispersion in propylene carbonate (the amount given is relative to 100 parts of the other components).

The components given in Table 1 were placed in a 100 ml beaker and kept under stirring, at room temperature, for 1 hour. They were then left to stand overnight in order to obtain a homogeneous composition free of bubbles.

#### EXAMPLE 6

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#### Mechanical and chemical-physical analyses

The compositions of Examples 1-5 were subjected to the following mechanical and chemical-physical analyses.

<u>Viscosity</u>

obtained according to Examples 1-5 was measured, at 30°C and at 80°C, using a viscometer of Brookfield type, model DV-III, equipped with a configuration 29. The results obtained are given in Table 2.

#### Modulus of elasticity values

Films were obtained from the abovementioned compositions by working as follows. A film 70  $\mu$ m in thickness and 120 mm in width was spread onto a glass plate using the "Bird" filmograph at a speed of 2 m per minute; the crosslinking of the film was carried out using a Fusion UV curing System device, model F600 and lamp with spectrum H, applying a UV dose of 1.25 J/cm². At the end of the crosslinking, the films were removed

1.00

from the glass plate.

The film obtained from the composition of Example 2 was not subjected to further analyses since said composition was found to have undergone little crosslinking; as a matter of fact, said composition had a sticky appearance and left residues on the surface of the glass at the time of removal.

The films thus obtained were conditioned for 24 hours, at 25°C and at 50% relative humidity, and were then subjected to measurement of the modulus of elasticity by means of a DMTA (Dynamic Mechanical Thermal Analyser from Reometrics Inc.), in traction, at a frequency of 1 Hz and at a heating rate of 2°C/min over the temperature range between -60°C and 120°C.

The results obtained, relating to the modulus values measured at room temperature (20°C) and at -40°C, are given in Table 2.

### H<sub>2</sub>O absorption of the crosslinked films

The films obtained as described above, predried in an atmosphere flushed with dry air for 48 hours, were subjected to controlled absorption of H<sub>2</sub>O. To this end, the Igasorp machine from Hiden Analytical was used, working at a temperature of 55°C, with a relacive humidity of 95%, until an asintotic value in the absorption of the water content was reached.

The results obtained, expressed as a percentage absorption of absorbed water, are given in Table 2.

Thermal ageing

The films obtained as described above were subjected to ageing for 8 days, at 80°C. The reduction in mechanical characteristics, in particular the elongation at break and the stress at break were then evaluated: the results obtained (the percentage variation is reported) are given in Table 2.

To this end, the mechanical characteristics were measured using an INSTRON 4502, Series 9 dynamometer, at

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a traction speed of 25 mm/min, on punches 150 mm in height and 20 mm in width obtained from the abovementioned films preconditioned at 25°C, with a humidity of 50%, for 24 hours. For comparative purposes, the mechanical characteristics were also measured on punches obtained from non-aged films.

The percentage variation in the mechanical characteristics was calculated relative to the value of said characteristics measured on punches obtained from the comparative (non-aged) films.

### Amount (%) of extractable materials after crosslinking

The amount of extractable materials was measured as follows. The films obtained as described above were immersed in distilled water contained in 250 ml beakers, said beakers were covered so as to limit the evaporation of the water and were then placed in an oven thermostatically maintained at 60°C. The treatment was continued for 15 days, filling up, if necessary, with distilled water when the level decreased. At the end, the extracted material (E) was calculated according to the following formula:

$$E = \frac{W_0 - W_1}{W_0} * 1.00$$

in which:

- $W_0$  represents the weight of the original film dried 25 at 60°C for 24 hours;
  - $W_1$  represents the weight of the film subjected to the abovementioned treatment and then dried at 60°C for 24 hours.

The results obtained are given in Table 2.

TABLE 2

	COMPOSITIONS					
	1 (*)	2 (*)	3	4	5	
Viscosity at 30°C	12.5	80.0	445	650	505	
(poise)						
Viscosity at 80°C	-	_	19.0	27.0	20.8	
(poise)		tomorphy and		Sec. 2.	egran i see	
Modulus at 20°C (MPa)	102	_	1.3	2.7	2.1	
Modulus at -40°C (MPa)	2540	_	<sup>:</sup> 36	30	34	
H <sub>2</sub> O absorption (%)	2.3	-	0.45	0.70	0.50	
Elongation at break	-94%	-	+6%	-15	+10	
(% variation)		141	संग्रेशकर <u>के</u> में के	, og e samples		
Stress at break (% variation)	+414%	-	+21%	+60	+39	
Extractable materials (%)	0.9		0.1	0.1	0.1	

(\*): comparative

The data given in Table 2 show that crosslinkable composition according to 5 the present invention (Examples 3, 4 and 5) is better than the comparative composition (Examples 1 and 2). In particular, the crosslinkable composition according to the present invention shows:

- lower modulus values at low temperatures;
- less water absorption;
- less variation in the elongation at break and in the stress at break;
- 5 smaller amount of extractable materials.

#### EXAMPLES 7-11

Preparation of compositions for primary coating with adhesion promoter

Compositions for primary coating with adhesion promoter according to the invention were prepared: the amounts of the components (parts by weight except where otherwise mentioned) are given in Table 3.

TABLE 3

-						
	COMPOSITIONS					
COMPONENTS	7	8	9	10	11	
(a) Kraton® Liquid	50	50	50	50	50	
L-207						
(b) Kraton® Liquid	50	50	50	50	50	
L-1203						
вүк® 361	0.5	0.5	0.5	0.5	0.5	
Silquest® A-187	-	1.0	_	<del>-</del>	-	
Silquest® A-186	<b>-</b> .	_	1.0	-		
Dynasylan® MTMO	-		-	1.0	_	
Si® 266					1.0	
UVI® 6974	0.5	0.5	0.5	0.5	0.5	
Timid Exp 207. linear eligemer containing a						

15 Kraton Liquid™ EKP-207: linear oligomer containing a poly(ethylene/butylene) aliphatic chain, a hydroxyl group at one end and epoxide groups at the other end, sold by Kraton Polymer;

Kraton Liquid™ L-1203: hydroxy-terminated hydrogenated

polydiene oligomer sold by Kraton Polymer;

BYK®-361: polyacrylate copolymer sold by BYK-Chemie [the amount given is relative to 100 parts of the components (a) + (b)];

- 5 Silquest® A-187: gamma-glycidoxypropyltrimethoxysilane sold by OSi Specialties [the amount given is relative to 100 parts of the components (a) + (b)];
  - Silquest® A-186: beta(3,4-epoxycycloexyl)ethyl-trimethoxysilane sold by OSi Specialties [the amount
- given is relative to 100 parts of the components (a) + (b)];

Dynasylan® MTMO: gamma-mercaptopropyltrimethoxysilane sold by Degussa-Hüls [the amount given is relative to 100 parts of the components (a) + (b)];

- 15 Si® 266: bis(3-triethoxysilylpropyl)disulfane sold by Degussa-Hüls [the amount given is relative to 100 parts of the components (a) + (b)];
  - UVI® 6974: hexafluoroantimony triarylsulphonium salt, a photo-initiator sold by Union Carbide, as a 50%
- 20 dispersion in propylene carbonate (the amount given is relative to 100 parts of the other components).

The components given in Table 3 were placed in a 100 ml beaker and kept under stirring, at room temperature, for 1 hour. They were then left to stand overnight in order to obtain a homogeneous composition free of bubbles.

#### Example 12

### Preparation of a composition for secondary coating

A composition for the secondary coating was 30 prepared: the amounts of the components (parts by weight except where otherwise mentioned) are given in Table 4.

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#### TABLE 4

	COMPOSITION
COMPONENTS	
(1) Poly Bd® 605	50
(2) Cyracure UVR® 6105	40
(3) Cyracure UVR® 6000	10
(4) UVI® 6974	1.5
(5) BIK® 361	0.5

Poly Bd® 605: epoxidized polybutadiene sold by Elf

5 Atochem;

Cyracure® UVR-6125: 3,4-epoxycyclohexylmethyl-3,4-epoxycyclohexylcarboxylate sold by Union Carbide;

Cyracure® UVR-6000: 3-ethyl-3-(hydroxymethyl)oxetane sold by Union Carbide;

- 10 UVI® 6974: hexafluoroantimony triarylsulphonium salt, photo-initiator sold by Union Carbice as a 50% dispersion in propylene carbonate [the amount given is relative to 100 parts of the components (1) + (2) + (3)];
- 15 BYK®-361: polyacrylate copolymer sold by BYK-Chemie [the amount given is relative to 100 parts of the components (1) + (2) + (3)].

The components given in Table 4 were placed in a 100 ml beaker and were kept under stirring for 1 hour at room temperature. They were then left to stand overnight so as to obtain a homogeneous composition free of bubbles.

The composition obtained was subjected to the following analyses: viscosity and modulus of elasticity

values, working as described above in Example 1. The results obtained are given in Table 5.

#### TABLE 5

•	COMPOSITION
Viscosity at 26°C	20.1
(poise)	
Modulus at 20°C	2010
(MPa)	
Modulus at -40°C	2760
(MPa)	

EXAMPLES 13

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#### Glass adhesion measurement

The compositions of Examples 7-12 were subjected to the following analysis.

10 Glass plates coated with films made from the compositions of Examples 7 to 11 (primary coating) and from the composition of Example 12 (secondary coating) were obtained by working as follows. A film of primary coating (70  $\mu\mathrm{m}$  thick and 100 mm width) was spread onto a glass plate previously conditioned at 130°C for 10 hours 15 using the "Bird" filmograph at a speed of 1 m per minute; the crosslinking of the film was carried out using a Fusion UV curing System device, model F600 and lamp with spectrum H, applying a UV dose of 1.5 J/cm2. At the end of the crosslinking, a composition for secondary 20 coating prepared according to Example 12, was spread as a film (70  $\mu\text{m}$  thick and 120 mm width) onto the said

cross-limked film and was subjected to the crosslinking operating at the working conditions above disclosed.

The glass plates thus obtained were conditioned for 24 hours, at 25°C and at 50% relative humidity, and were subsequently immersed in distilled water for 24 hours at (aged samples). Αt the end of said treatment, the glass plates were subjected to adhesion measurement. To this end, from the mentioned glass plates samples having about 140  $\mu$ m in thickness and 40 mm in width were obtained. A traction 10 force in a direction perpendicular to the glass surface at a traction speed of 50 mm/min was applied to said samples, using an INSTRON 4502, Series 9 dynamometer equipped with a movable platform and a load cell of 10 N. In order to work in water saturation conditions, said 15 measurements were carried out not more then 10 minutes after the glass plates were extracted from the water. For comparative purposes, the glass adhesion measurement was also carried out on samples obtained from non-aged 20 films.

The results obtained, expressed in Newton/meter (N/m), are given in Table 6.

TABLE 6 - ----

•		
COMPOSITIONS	NON-AGED FILMS	AGED FILMS
	(N/m)	(N/m)
7	62.5	5.0
8	67.3	32.5
9	84.3	21.2
10	113.3	58.0
11	68.1	83.9

The data given in Table 6 show that the addition of an adhesion promoter to the crosslinkable composition according to the present invention improve the adhesion between the glass fiber and the primary coating, in particular upon ageing of the coating.

#### EXAMPLE 14

#### Production of optical fibers

Two optical fibers were produced according to the techniques known in the art, comprising a primary coating according to the present invention (compositions of Examples 7 and 10) and a secondary coating prepared according to Example 12. The primary coating was applied at a temperature of 80°C as the secondary coating was applied at a temperature of 26°C. The spinning speed was 14 m/s. The primary coating and the secondary coating were applied to a thickness of 30  $\mu$ m each.

#### EXAMPLE 15

#### Strip test

Two test specimens of the optical fibers obtained as

20 disclosed in Example 14 were subjected to a strip test
according to the Bellcore standard GR-20-CORE, July
1998, section 4.4.2. To this end, the specimens were
conditioned at room temperature, with a humidity of 50%,
for 7 days and subsequently were immersed in water at

25 20°C and at 60°C for 7 days (aged samples). For
comparative purposes, the strip test was also carried
out on samples obtained from non-aged specimens.

The result obtained, which are the average of 5 different tests, are given in Table 7.

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TABLE 7

STRIP FORCE VALUE	Example 7	Example 10
(N)		
NON-AGED	1.00	1.82
AGED	0.00	1.00
(7 days at 20°C)		
AGED	0.00	0.78
(7 days at 60°C)	40	

The data given in Table 7 show that the addition of an adhesion promoter to a crosslinkable composition 5 according to the present invention improve the adhesion between the glass fiber and the primary coating, particular upon ageing of the coating.

### EXAMPLE 16

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### Attenuation measurement

A 1 000 m skein of the optical fiber obtained as - many disclosed in Example 14 comprising a primary coating invention (composition of according to the present Example 7) and a secondary coating prepared according to Example 12, was subjected to attenuation measurements using an OTDR (optical time domain reflectometer) from ANRITSU, model MW 90-60 A.

The attenuation measurements carried out at 20°C and at 1550 nm gave a value of 0.20 dB/km, while the attenuation measurements carried out at 20°C and at 1330 nm gave a value of 0.35 dB/km. The use of the primary coating according to the present invention thus gives the optical fiber good attenuation characteristics.

## CLAIMS

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- 1. Optical fiber comprising at least one epoxidized polyolefin based polymer coating, characterized in that said coating is formed from a crosslinkable composition comprising:
  - (a) at least one epoxidized polydiene oligomer having a first and a second end, said oligomer comprising at least one hydrocarbon chain that is substantially free of ethylenic double bonds, at least one epoxide group at said first end and at least one reactive functional group at said second end;
- (b) at least one hydrogenated polydiene oligomer
  comprising at least one reactive functional group
  capable of reacting with said epoxide groups;
  - (c) at least one photo-initiator.
  - 2. Optical fiber according to Claim 1, in which said polymer coating is a primary coating.
- 3. Optical fiber according to Claim 1, in which the crosslinkable composition comprises at least one adhesion promoter (d).
  - 4. Optical fiber according to Claim 1, in which the crosslinkable composition comprises at least one reactive diluent monomer (e).
  - 5. Optical fiber according to Claim 1, in which the crosslinkable composition has a modulus of elasticity, at room temperature, of less than about 4 MPa.
- 6. Optical fiber according to Claim 5, in which the crosslinkable composition has a modulus of elasticity, at room temperature, of between 1 MPa and 3 MPa.
- 7. Optical fiber according to Claim 1, in which the crosslinkable composition has a modulus of elasticity, at -40°C, of between 5 MPa and 350 MPa.

- 8. Optical fiber according to Claim 7, in which the has а modulus composition crosslinkable elasticity, at -40°C, of between 10 MPa and 50 MPa.
- 9. Optical fiber according to Claim 1, in which the epoxidized polydiene oligomer (a) is obtained by 5 of conjugated (co)polymerization monomers containing from 4 to 24 carbon atoms, or of disubstituted conjugated diene monomers, or difluorinated conjugated diene monomers, optionally copolymerized with other ethylenically unsaturated 10 followed by functionalization, monomers, hydrogenation and epoxidation.
- 10. Optical fiber according to Claim 9, in which the epoxidized polydiene oligomer (a) is a copolymer comprising a first block comprising at 15 epoxide group, obtained by the least one polymerization of a first conjugated diene monomer, which is subsequently epoxidized, and a second block hydrocarbon chain that from a formed ethylenic double 20 substantially free of polymerization of obtained by the conjugated diene monomer, subsequently terminated with at least one reactive functional group and hydrogenated.
- 11. Optical fiber according to Claim 10, in which the 25 first conjugated diene monomer is isoprene.
  - 12. Optical fiber according to Claim 10, in which the second conjugated diene monomer is 1,3-butadiene.
- 13. Optical fiber according to Claim 1, in which from 5 to 15 epoxide groups are present in the epoxidized 30 polydiene oligomer (a), at the first end.
  - 14. Optical fiber according to Claim 13, in which from 9 to 11 epoxide groups are present in the epoxidized polydiene oligomer (a), at the first end.
- 15. Optical fiber according to Claim 1, in which, in the 35 epoxidized polydiene oligomer (a), the reactive

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- functional group present at the second end is selected from: hydroxyl, acrylate, epoxy, vinyl ether, mercaptan.
- 16. Optical fiber according to Claim 15, in which, in the epoxidized polydiene oligomer (a), the reactive functional group present at the second end is a hydroxyl group.
- 17. Optical fiber according to Claim 1, in which the epoxidized polydiene oligomer (a) has an average molecular weight of between 3,000 daltons and 15,000 daltons.
  - 18. Optical fiber according to Claim 1, in which the epoxidized polydiene oligomer (a) has a viscosity, measured at 30°C, of less than 1,000 poise.
- 19. Optical fiber according to Claim 1, in which the hydrogenated polydiene oligomer (b) has a base polymer structure which is derived from the (co)polymerization of one or more conjugated diene monomers containing from 4 to 24 carbon atoms, optionally copolymerized with other ethylenically

unsaturated monomers.

- 20. Optical fiber according to Claim 1, in which, in the hydrogenated polydiene oligomer (b), the reactive functional group is selected from: hydroxyl, acrylate, epoxy, vinyl ether, mercaptan.
- 21. Optical fiber according to Claim 20, in which, in the hydrogenated polydiene oligomer (b), the reactive functional group is a hydroxyl group.
- 22. Optical fiber according to Claim 21, in which, in the hydrogenated polydiene oligomer (b), the hydroxyl group is in an end position.
  - 23. Optical fiber according to Claim 22, in which the hydrogenated polydiene oligomer (b) has a hydroxyl functionality of between about 0.5 and about 2.6.
- 35 24. Optical fiber according to Claim 1, in which the hydrogenated polydiene oligomer (b) has a viscosity,

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- measured at 30°C, of between about 10 poise and about 1,000 poise.
- 25. Optical fiber according to Claim 1, in which the hydrogenated polydiene oligomer (b) has an average molecular weight of between about 500 daltons and about 20,000 daltons.
- 26. Optical fiber according to Claim 1, in which the hydrogenated polydiene oligomer (b) has a hydroxyl-250 and equivalent weight of between about about 20,000.
- 27. Optical fiber according to Claim 1, in which the from: selected (c) is photo-initiator salts, triarylsulphonium hexafluorophosphorus triarylsulphonium salts, hexafluoroantimony
- (tolylcumy1) tetrakis (pentafluoropheny1) iodonium 15 salts, diaryliodonium hexafluoroantimonate salts, or mixtures thereof.
  - 28. Optical fiber according to Claim 3, in which the adhesion promoter is selected from organo-functional silanes.
- 29. Optical fiber according to Claim 28, in which the gammais organo-functional silane glycidoxypropyltrimethoxysilane, beta (3,4 epoxycycloexhyl)ethyltrimethoxysilane, gammamercaptopropyltrimethoxysilane.
  - 30. Optical fiber according to Claim 29, in which the gammais silane compound organo-functional mercaptopropyltrimethoxysilane.
  - 31. Optical fiber according to claim 28, in which the following the organo-functional silane has 30 structural formula (I):

#### $(R)_3Si-C_nH_{2n}-X$ (I)

in which the groups R, which may be identical to or different from each other, are selected from: alkyl, alkoxy or aryloxy groups or from halogen atoms, on condition that at least one of the groups R is an

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- alkoxy or aryloxy group; n is an integer between 1 and 6 inclusive; X is a group chosen from: nitrous, mercapto, epoxide, vinyl, imido, chloro,  $-(S)_mC_nH_{2n}-Si-(R)_3$  in which m and n are integers between 1 and 6 inclusive and the groups R are defined as above.
- 32. Optical fiber according to Claim 31, in which the organo-functional silane compound is bis(3-trimethoxysilylpropyl)disulfane, bis(3-
- 10 triethoxysilylpropyl)disulfane.
  - 33. Optical fiber according to Claim 3, in which the adhesion promoter is added to the crosslinkable composition in an amount of from 0.1 parts by weight to 2.5 parts by weight relative to 100 parts of (a) + (b).
  - 34. Optical fiber according to Claims 33, wherein the adhesion promoter is added to the crosslinkable composition in an amount of from 0.3 parts by weight to 1.5 parts by weight relative to 100 parts of (a) + (b).
  - 35. Optical fiber according to Claim 4, in which the reactive diluent monomer is selected from vinyl ethers.
- 36. Optical fiber according to Claim 4, in which the
  amount of reactive diluent monomer added to the
  crosslinkable composition is not greater than 20
  parts by weight relative to 100 parts of (a) + (b).
  - 37. Crosslinkable composition comprising:
- (a) at least one epoxidized polydiene oligomer having
  a first and a second end, said oligomer
  comprising at least one hydrocarbon chain that is
  substantially free of ethylenic double bonds, at
  least one epoxide group at said first end and at
  least one reactive functional group at said
  second end;
  - (b) at least one hydrogenated polydiene oligomer

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comprising at least one reactive functional group capable of reacting with said epoxide groups;

- (c) at least one photo-initiator.
- 38. Crosslinkable composition according to Claim 37,5 comprising:
  - (a) about 20-80 parts by weight of at least one epoxidized polydiene oligomer having a first and a second end, said oligomer comprising at least one hydrocarbon chain that is substantially free of ethylenic double bonds, at least one epoxide group at said first end and at least one reactive functional group at said second end;
  - (b) about 20-80 parts by weight of at least one hydrogenated polydiene oligomer comprising at least one reactive functional group capable of reacting with said epoxide groups;
  - (c) about 0.05-5 parts by weight relative to 100 parts of (a) f (b) of a photo-initiator.
- 39. Crosslinkable composition according to Claim 37 or 38, in which the epoxidized polydiene oligomer (a) is as defined in any one of Claims 9 to 18.
  - 40. Crosslinkable composition according to Claim 37 or 38, in which the hydrogenated polydiene oligomer (p) is as defined in any one of Claims 19 to 26.
- 25 41. Crosslinkable composition according to Claim 37 or 38, in which the photo-initiator (c) is as defined in Claim 27.
  - 42. Crosslinkable composition according to Claim 37 or 38, in which at least one adhesion promoter (d) is present.
    - 43. Crosslinkable composition according to Claim 42, in which the adhesion promoter is defined in any one of Claims 28 to 34.
- 44. Crosslinkable composition according to Claim 37 or 38, in which at least one reactive diluent monomer (e) is present.

- 45. Crosslinkable composition according to Claim 44, in which the reactive diluent monomer is as defined in Claims 35 or 36.
- 46. Method for applying an epoxidized polyolefin based coating to an optical fiber, which comprises:
  - drawing a glass preform placed in a suitable furnace;
  - cooling the fiber leaving the furnace;
  - applying said coating;
- crosslinking said coating;

  characterized in that the application of said coating layer is carried out at a temperature of not less than 60°C.
- 47. Method according to Claim 46, in which the coating
  15 layer is formed from a crosslinkable composition
  defined according to any one of Claims 37 to 45.

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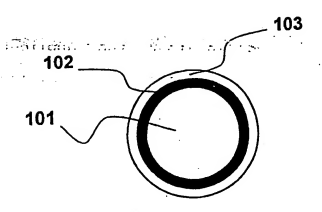


Fig. 1

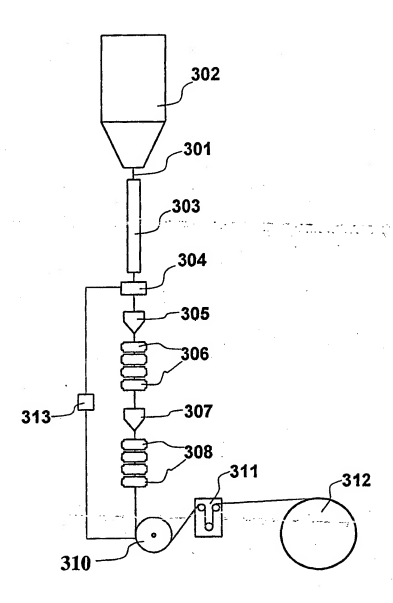


Fig. 2

## INTERNATIONAL SEARCH REPORT

Internation Application No PCT/EP 02/13447

A. CLASSI	FICATION OF	SUBJECT	MATTER
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According to International Patent Classification (IPC) or to both national classification and IPC

#### B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)  $IPC\ 7\ CO3C$ 

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, PAJ, WPI Data

C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		,
Category *	Citation of document, with indication, where appropriate, of	the relevant passages	Relevant to claim No.
X	US 6 042 943 A (LEVY ALVIN C) 28 March 2000 (2000-03-28) column 6 column 10 claims 1-80; examples		1-46
Х	EP 0 516 203 A (SHELL INT RESEARCH) 2 December 1992 (1992–12–02) example 5		1
A	EP 0 533 397 A (AMERICAN TELE TELEGRAPH) 24 March 1993 (199 the whole document		1-46
V Furt	her documents are listed in the continuation of box C.	Patent family members are listed	In anney
X Furi	tiel documents are used in the continuation of box ().	X Faterit rating members are insert	III cutilex.
'A' docume consider a docume which citation 'O' docume other a 'P' docume'	and defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another n or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but han the priority date claimed	"T" later document published after the Inter or priority date and not in conflict with cited to understand the principle or the invention.  "X" document of particular relevance; the cannot be considered novel or cannot involve an inventive step when the document of particular relevance; the cannot be considered to involve an indocument is combined with one or ments, such combination being obvious in the art.  "&" document member of the same patent.	the application but early underlying the claimed invention to considered to coument is taken atone claimed invention early step when the one other such docu-us to a person skilled
Date of the	actual completion of the international search	Date of malling of the international se	arch report
6	February 2003	13/02/2003	
Name and I	mailing address of the ISA	Authorized officer	
	European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Marquis, D	

# INTERNATIONAL SEARCH REPORT

Internate Application No
PCT/EP 02/13447

Category °	ation) DOCUMENTS CONSIDERED TO BE RELEVANT  Citation of document, with Indication, where appropriate, of the relevant passages		Relevant to old-
A	PATENT ABSTRACTS OF JAPAN vol. 004, no. 101 (C-019), 19 July 1980 (1980-07-19) -& JP 55 065242 A (MITSUBISHI CHEM IND LTD), 16 May 1980 (1980-05-16) abstract		Relevant to claim No.
A	US 5 993 965 A (FLAT JEAN-JACQUES ET AL) 30 November 1999 (1999-11-30) the whole document	*	1–46
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## INTERNATIONAL SEARCH REPORT

information on patent family members

Internati Application No PCT/EP 02/13447

<del> </del>		·			
Patent document cited in search report		Publication date		Patent family member(s)	Publication date
US 6042943	Α _	28-03-2000	US	5985952 A	16-11-1999
			WO	02096180 A1	05-12-2002
EP 0516203	Α	02-12-1992	US	5229464 A	20-07-1993
		•	US	5382604 A	17-01-1995
			BR	9201539 A	01-12-1992
			CN	1066275 A ,B	18-11-1992
		•	CN	1244550 A	16-02-2000
			DE	69232787 D1	31-10-2002
		•	EP	0516203 A2	02-12-1992
			JP	2870622 B2	17-03-1999
			JP	5125197 A	21-05-1993
			KR	227998 B1	01-11-1999
	(		MX	9201967 A1	01-11-1992
			RU	2101295 C1	10-01-1998
			US	5399626 A	21-03-1995
•			ÜS	5449718 A	12-09-1995
			US	5686535 A	11-11-1997
•			US	5389701 A	14-02-1995
			US	5491193 A	13-02-1996
ED 052227		04 00 1000			
EP 0533397	Α	24-03-1993	US	5181269 A	19-01-1993
			AU	654209 B2	27-10-1994
			AU	2456192 A	18-03-1993
			CA	2074909 A1	18-03-1993
			CN	1070738 A ,B	07-04-1993
			EP	0533397 A1	24-03-1993
			JP	2690433 B2	10-12-1997
			JP	6250053 A	09-09-1994
			MX	9205227 A1	01-03-1993
JP 55065242	Α	16-05-1980-	JP	1374836 C	22-04-1987
			JP	61043376 B	27-09-1986
·					
US 5993965	Α	30-11-1999	AU	744485 B2	28-02-2002
en skip eerskip in groot	richerter in	·	AU	5467898 A	20-08-1998
• • • • • • • • • • • • • • • • • • •	•	•	BR	9800637 A	14-12-1999
			CA	2229974 A1	18-08-1998
•		•	CN	1196419 A ,B	21-10-1998
			EP	0859038 A1	19-08-1998
			JP	10232331 A	

